PATENT SPECIFICATION



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Inventors: LESLIE ALFRED MUIRHEAD, EDWARD LAWRENCE HILL and BRIAN WILFRED FORSTER.

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COMPLETE SPECIFICATION

DRAWINGS ATTACHED

Improvements in and relating to the Manufacture of Particulate Expandable Polymers and to the Manufacture of Expanded Polymers Therefrom

We, SHELL INTERNATIONALE RESEARCH MAATSCHAPPIJ N.V., a company organised under the laws of The Netherlands, of 30 Carel van Bylandtlaan, The Hague, The 5 Netherlands, do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement:

The present invention is concerned with the manufacture of expandable particles of a polymer of a vinyl-aromatic compound and for a vinylidene-aromatic compound (hereinafter referred to as a thermoplastic organic 15 polymer), for example, polystyrene the term polymer including a copolymer or a polymer composition; with the resulting expandable particles; and with the manufacture of an expanded (i.e. a foamed) polymer of cellular 20 form, including shaped articles, therefrom. It is known that a thermoplastic organic

polymer such as polystyrene can be expanded to a cellular form by intimately incorporating a vaporisable expanding agent 25 in the polymer and then heating the polymer under conditions permitting the expanding agent to vaporise to form a larger number of individual, enclosed spaces or cells within the In the case of polystyrene it is polymer. 30 common to use as the vaporisable expand-

ing agent a normally-liquid, vaporisable

organic substance, such as technical pentane, which when intimately incorporated in the polymer forms an "expandable polymer."

35 On heating such an expandable polymer, the expanding agent vaporises in a large number of individual locations within the polymer to form said cells. On cooling, the resulting polymer is of cellular form: such a polymer 40 is referred to herein as an "expanded poly-

mer.

It is also known to manufacture shaped, cellular articles from polymers by forming

an expandable polymer in particulate form, i.e. expandable polymer particles, and then 45 heating a mass of "pre-expanded" expandable particles in an enclosed mould of appropriate shape. The expandable polymer particles are normally "pre-expanded", that is they are heated to effect at least a major 50 part of the expansion of which they are capable, prior to charging the particles to the mould so that formation of the shaped cellular article is in fact effected by the fusion together of the pre-expanded particles in the 55 mould with only a minor amount of further expansion occurring therein. An early description of such technique occurs in U.K. Patent Specification No. 715,100. To avoid the necessity for transporting the more bulky 60 pre-expanded particles from the manufacturer of expandable particles to the user thereof, pre-expansion of expandable par-ticles is usually carried out by the user, i.e. the manufacturer of shaped cellular poly- 65 mer articles, and such pre-expansion is usually effected by contacting the expandable polymer particles with live steam.

For many years expandable polystyrene particles have been manufactured by incor- 70 porating a vaporisable expanding agent in polystyrene beads obtained by a suspension polymerisation process. More recently it has also been proposed to incorporate such an expanding agent in heat-plastified poly- 75 styrene in an extruder and to form expandable particles by cutting the expandable, filamentary extrudate. Since such an extrudate is an expandable material at an elevated temperature the prior proposals have in-80 cluded special steps designed to prevent any substantial premature expansion of the extrudate, such steps comprising immediately contacting the extrudate with a liquid coolant such as water to cool the extrudate to 85 a temperature below 50°C or extruding the

expandable polymer into a pressure zone in which the extrudate remains until it has been cooled to a temperature at which it is incapable of expanding to any significant extent.

The use of a pressure zone brings with it operational complications, particularly when the extrudate is chopped into particles within the pressure zone; whilst sudden cooling of the extrudate to below 50°C is said to "freeze" the polymer in an oriented or strained condition and in such case a subsequent annealing operation involving heat treatment at a temperature between 50 and 75°C has been proposed as a means of relax-

15 ing the orientation or strain existing within the expandable polymer particles.

According to the present invention a process for the manufacture of expandable particles of a thermoplastic organic polymer 20 (as hereinbefore defined) in which a vaporisable expanding agent is incorporated in a heat-plastified mass of said polymer and the heat-plastified polymer is extruded in filamentary form and cut into particles, which 25 comprises extruding the heat-plastified polymer, containing said expanding agent, into an essentially gaseous environment at atmos-

pheric or substantially atmospheric pressure and immediately cooling the extruded poly-30 mer by contact with a fluid at a temperature between 60 and 80°C in such a manner that the temperature of the extruded polymer does not fall below 60°C, maintaining the temperature of the extruded polymer be-

temperature of the extruded polymer be-35 tween 60 and 80°C during a normalising period (as herein defined) immediately following extrusion thereof and before and/or after cutting the extruded polymer into particles, and thereafter cooling the polymer 40 particles to ambient temperature. Prefer-

ably the extruded polymer is cut into particles and the resulting particles are then subjected to normalising conditions, although it is also possible to normalise the 45 polymer in filamentary form prior to cutting

or for the normalising period to extend both before and after the cutting operation.

The present invention also includes the resulting particulate expandable polymer i.e. 50 expandable particles of a thermoplastic organic polymer; and expanded particles and shaped articles formed therefrom by any known or suitable expansion technique or moulding process.

The reference herein to an essentially gaseous environment means an environment which is either wholly gaseous or mainly gaseous to the extent that liquid droplets, for example, a water spray can also be present.

example, a water spray can also be present.

60 Said normalising period is defined herein as the minimum period, i.e. length of time, for which an extruded expandable polymer in filamentary or particulate form must be maintained at a temperature within the range 65 60 to 80°C and at normal pressure prior to

being cooled to ambient temperature in order that the size of at least 90% of the cells of an expanded particle, obtained by freely expanding such an expandable particle (or an expandable particle cut from such expandable flamentary polymer) by heating under conditions resulting in a maximum increase in volume of the particle, is not greater than 100 microns (in diameter). In general a cell size of about 100 microns is the maximum cell size currently acceptable commercially in the manufacture of cellular polystyrene articles by moulding expanded polystyrene particles.

The process of the present invention has 80 the operational advantage of producing normalised expandable polymer particles which do not require any subsequent annealing operation and since the temperature of the extruded polymer is not allowed to fall be- 85 low 60°C until normalisation has been achieved, it is possible that one avoids the intermediate stage in which, according to the prior art, orientation or strain is produced. It has been found that operation in accord- 90 ance with the present invention permits the attainment of expandable polystyrene particles which, from the point of view of their end-use behaviour in a subsequent moulding operation leading to shaped cellular poly- 95 styrene articles, closely resemble the wellknown expandable polystyrene particles obtained by incorporating an expanding agent in the particulate product of a suspension

polymerisation process.

One of the major difficulties which can arise in the manufacture of expandable polymer particles by methods involving incorporating a vaporisable expanding agent in the heat-plastified polymer is the problem of 105 cell size in the resulting expanded polymer and, whatever the mechanism of the normalising treatment of the present invention may be, it has been found that operation in accordance with the present invention enables a satisfactory cell size (i.e. 90% below 100 microns) to be attained in the ultimate

cellular polymer.

In carrying out the present invention it is essential to normalise the extruded expandable polymer either as particles or in filamentary form prior to cutting into particles. The extrudate comprising a thermoplastic organic polymer, for example, polystyrene in a heat-plastified form and containing a responsable expanding agent dispersed therein, can be cooled to and normalised at 60-80°C whilst in the form of a filament (or a plurality of filaments). Such a normalised filament can then be cut into particles whilst said filament is at 60-80°C or after first cooling to ambient temperature; but preferably said filamentary extrudate is cut into particles immediately following extrusion thereof by cutting the extrudate at the extrusion ori-

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fice of the extruder (face-cutting) or shortly after extrusion so that the extruded expandable polymer is normalised in particulate Operation in this preferred manner 5 has practical advantages as compared to normalisation in filamentary form. When employing the preferred technique the extruded filament is immediately contacted, in an atmospheric or substantially atmospheric pressure 10 environment, with a fluid at a temperature between 60 and 80°C whilst simultaneously, or within a short time of extrusion, cutting said extruded filament into particles, and thereafter contacting the resulting particles 15 with a fluid at a temperature between 60 and 80°C under conditions such that the temperature of both said extruded filament and the resulting particles does not fall below 60°C during the required normalising period 20 immediately following extrusion.

The extruded filament can be cooled by contact with a liquid or gas at a temperature of 60-80°C, but preferably with a liquid, e.g. water. When face-cutting is employed the liquid is preferably in droplet form, the expandable polymer being sprayed therewith as it is being extruded. When the filamentary extrudate is cut into particles subsequent to cooling with a liquid at 60-80°C. 30 the extrudate can be passed immediately into a body of such liquid disposed as near as possible to the extrusion orifice(s) and if desired the extrudate can also be sprayed with liquid prior to entering the body of liquid. 35 The normalising of the extruded filament or ticles with a body of cooling liquid at 60-80°C, which is preferably water. Cooling 40 of the normalised filament(s) or particles to ambient temperature is conveniently achieved by contacting said filament(s) or particles with air and this can be a combined cooling and drying operation to re-

therewith. Whilst the temperature of the heat-plasti-fied polymer as it is being extruded (i.e. its extrusion temperature at the die-head) will 50 depend, inter alia, on the softening tem-perature of the polymer (which may be lower than that of the polymer per se due to the presence therein of the vaporisable expanding agent), the extrusion temperature should 55 always be high enough to avoid the possi-bility of solidification of the polymer in the extrusion orifice(s) of the die-head. required extrusion temperature will usually depend on the dimensions, and in particular 60 the cross-sectional area, of the extrusion orifice; and on the extent to which the die-head itself is cooled by the cooling fluid at 60-80°C with which the extrudate is contacted, particularly when using face-cutting tech-65 nique. For example, for polystyrene con-

45 move liquid, e.g. water, physically associated

taining 4 to 8% by weight of technical pentane as the expanding agent, the extrusion temperature of the heat-plastified polymer can be as low as between 105 and 115°C e.g. about 110°C, if the extrusion orifice(s) is 70 (are) of the order of 2 to 3 millimetres in diameter and if there is no significant heat loss from the heat-plastified polystyrene in the die-head, but higher temperatures should be employed when extrusion orifices of 75 smaller diameter are employed. In general extrusion temperatures for polystyrene containing technical pentane can be up to 150 to 160°C depending on the extrusion conditions employed. Temperatures of 130 to 80 160°C are advantageous when using small diameter extrusion orifices i.e. below 1.0 millimetres, e.g. of the order of 0.5 to 0.75 millimetres, in diameter.

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Normalising can be carried out by contacting the extrudate with said fluid at a temperature between 60 and 80°C e.g. about 70°C for a period, e.g. between 30 and 60 minutes for expandable polystyrene particles, depending on the physical form, e.g. the 90 particle size, of the extrudate. A normalising liquid, for example, water or brine or other aqueous media, e.g. mixtures of water and glycerol or ethylene glycol, can be used, and when the extrudate is normalised in 95 particulate form the particles are subsequently separated from the normalising liquid in any convenient manner, e.g. by employing a screen which can be an inclined, vibratable screen or a perforated travelling 100 belt. When a normalising liquid such as water is used the particles are thereafter dried and cooled to ambient temperature e.g. in a current of air.

It will be understood that a plurality, e.g. 105 between 20 and 50, of filaments of expandable polymer will usually be extruded simultaneously using a multi-orifice die-head when carrying out the present invention on a manufacturing scale. It is usual to employ circular extrusion orifices with the result that the filamentary extrudate is of essentially circular cross-section: whilst such a filament is of rod-like form it is usual in the art to refer to it as a strand and the term 115 "strand" will be used in this general sense in the following specific description.

Said thermoplastic polymer can be polystyrene or other polyvinyl-aromatic compound and/or polyvinylidene-aromatic compound or a copolymer of a vinyl-aromatic compound and/or vinylidene-aromatic compound e.g. with acrylonitrile, methyl methacrylate, acrylic acid or methacrylic acid; although the present invention is particularly applicable to polystyrene.

Said vaporisable expanding agent can be

Said vaporisable expanding agent can be a normally liquid organic substance having substantially no solvent action on said thermoplastic polymer or a mixture of such 130

organic substances; and if desired such a substance or mixture thereof can be used together with a small amount of a solvent for said polymer. Preferably said expanding agent is an organic liquid boiling, at normal pressure, below 95°C and usually below about 80°C, e.g. a hydrocarbon or mixture thereof such for example as n-pentane or technical pentane or a mixture of n-pentane 10 and iso-pentane containing between 30 and 80% by weight of iso-pentane. If desired said polymer can contain an additive, e.g. a low molecular weight polyethylene wax, advantageously in an amount which is between 15 0.05 and 0.1% by weight of the polymer, as referred to in the specification of our copending U.K. Patent Application No. 146/62 Serial No. 997356. The process of the present invention can 20 be carried out either by cutting the expandable polymer as it is extruded from the die orifice(s) using any known or suitable facecutting technique or by extruding the polymer in strand form and subsequently chop-25 ping or cutting the strand(s) into particles using any known or suitable strand-cutting In each case the expandable technique. polymer is immediately contacted with cooling fluid at 60 to 80°C, either as it issues 30 from the die orifice(s) as when face-cutting or as soon as practicable thereafter when a strand cutting technique is employed. When strand-cutting is used it is preferable to carry out the cutting operation before or at 35 least during the initial stage of the normalising period in order to avoid handling long strands: and preferably the degree of drawdown of the extruded straud(s) is as small as possible to reduce the risk of strand break-40 age and in general the degree of drawdown should not exceed that which will normally occur in a strand-cutting operation employing the minimum haul-off tension in the When employing the strand-cutstrands. 45 ting technique it is usually unnecessary to direct cooling fluid against the face of the die-head providing the extruded strands are substantially immediately contacted with said cooling fluid so as to reduce the tem-50 perature of at least the surfaces of the strands to the normalising temperature of 60 to 80°C as quickly as possible. However, it is possible to direct cooling fluid, e.g. a spray of liquid droplets, against the face of the die-55 head if desired and operation in this way is recommended if the extrudate tends to expand prematurely prior to entering for example a body of cooling liquid at 60 to 80°C disposed adjacent the die-head. In general, 60 however, when using water at 60 to 80°C as the cooling fluid, no cooling of the diehead itself will occur when employing a strand-cutting technique, whereas when em-ploying a face-cutting technique it has been 65 found advisable to ensure that the cutting

face is kept "wetted" with a cooling liquid in order to prevent smearing of the polymer on the cutting face, which impairs the face-cutting operation. The removal of heat from the die-head as the result of cooling liquid 70 contacting the die face could cause premature solidification of polymer in the diehead. However, this can be prevented in several ways including, for example, supplying heat to the die-head itself by pro- 75 viding it with an electrical heating element or passage for the circula-tion therethrough of a heating fluid e.g. steam, preferably together with the provision of thermal insulation between the die face 80 and the body of the die-head as will be described later in detail in connection with the use of face-cutting technique in carrying out the present invention. Alternatively or in addition one can operate at higher extrusion 85 temperatures, e.g. for polystyrene between 130 and 160°C, providing, of course, the extrusion temperature is not so high as to cause undesirable expansion of the expandable polymer as it issues from the die-head. 90 Whilst such measures are considered to be more important in the case of face-cutting, it is to be understood that the relationship between polymer solidification and extrusion orifice dimensions referred to previously can 95 render such measures applicable to strand cutting as well.

Whether face-cutting or strand-cutting techniques are employed, the vaporisable expanding agent is incorporated in the poly- 100 mer in a screw extruder comprising a barrel providing a heat-plastifying section followed by a dispersion section to which the expanding agent is supplied and in which it is incorporated in the heat-plastified polymer fed 105 by the extruder screw from the first section. Heat is supplied to the polymer in the first section, e.g. electrically or by means of a heat-transfer fluid circulated through a jacket surrounding the first section; and usually 110 heat is removed from the polymer in the second section by means of a heat-transfer fluid e.g. water circulated through a jacket surrounding the second section. truder screw extends through the length of 115 both sections of the extruder and is profiled to provide for agitation and forward feeding of the polymer in the first section and the exertion of extrusion pressure thereon; and for mixing of the heat-plastified polymer and 120 the expanding agent in the second section. The second section is followed by the diehead through which the expandable polymer formed in the second section is extruded under pressure exerted on the polymer 125 in the first section, the degree of heat removal in the second section being controlled in accordance with the desired extrusion temperature.

A convenient way of carrying out the pre- 130

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sent invention employing a strand cutting technique for manufacturing e.g. expandable polystyrene particles is illustrated diagrammatically in Figure I of the accompanying 5 drawings. Referring to Figure I, polystyrene granules are fed to a screw extruder 10 having a heating jacket 11 extending over part of its length in which the extruder screw 12 is of conventional profile for heat-plastify-10 ing the polystyrene and advancing the heatplastified polymer under extrusion pressure towards the extruder die-head 13. The retowards the extruder die-head 13. maining part 14 of the extruder 10 constitutes a dispersion section in which tech-15 nical pentane supplied through a line 15 is incorporated in the heat-plastified polystyrene. A cooling jacket 16 surrounds the dispersion section and that part 17 of the extruder screw lying within the dispersion 20 section is shaped to provide the agitation required to effect an intimate dispersion of the pentane in the polystyrene. The pentanecontaining polystyrene is extruded as strands 18 from the die-head 14 (four such strands 25 being shown) and these strands 18 are immediately led into a bath 19 containing water at between 60°C and 80°C to effect cooling of at least the skin of the strands 18 to the temperature of the water before 30 cutting the strands into particles using a conventional strand cutter 20. The resulting particles 21 fall into a slurry tank 22 also containing water at between 60°C and 80°C which constitutes a reservoir for the sub-35 sequent normalising stage which is carried out in a tower 23 comprising an internal agitator 24 shaped in such a manner as to progressively lift up through the tower 23 the particles supplied to the base of the tower 40 as an aqueous slurry (suspension), via a line 25, a pump 26 and a line 27, from the slurry tank 22. The tower 23 also contains an internal heating coil 28 for maintaining the water in the tower 23 at a temperature be-45 tween 60 and 80°C. The size of the tower The size of the tower 23 in relation to the average residence time of the particles 21 in the slurry tank 22 and the particle throughput rate in the tower 23 is such that the particles formed by the cut-50 ter 20 are subjected to the required normalisation i.e. the particles are maintained at a temperature between 60 and 80°C for a normalising period (e.g. 40 minutes) prior to discharge from the top of the tower 23. The normalised particles are discharged from the tower 23 through a line 29 onto the lower end of a conveyor 30 having a perforated belt to permit water to drain from the particles into a reservoir 31 in which the water 60 is heated for return via line 32 to the slurry tank 22. The particles 21 are conveyed to a bin 33, if desired via an air dryer shown in dotted lines at 34. In the alternative way of carrying out the 65 process of the present invention in which the

extruded polymer containing a vapourisable, normally liquid expanding agent e.g. technical pentane, is extruded and face-cut at the die-head, the face-cutting is preferably effected by means of a rotatable cutter whilst 70 the face of the die-head is simultaneously contacted with droplets of water at a temperature of 60 to 80°C sprayed onto said The water droplets at this temperature are conveniently formed by spraying 75 wet steam towards the face of the die-head. The resulting particles of expanded polymer are washed by the water spray into a body of water maintained at a temperature of 60 to 80°C, e.g. about 70°C, in which the par-80 ticles remain for the desired normalising period of e.g. 30 to 60 minutes. The diehead is provided with means for counteracting the heat loss which occurs as the result of the water spray on the die face and 85 a preferred way of achieving this is to employ a die-head of sandwich construction which comprises a front plate constituting the die face against which the cutter rotates, a back plate which is secured to the extrusion 90 end of the extruder and, between the front and back plates, an insulating plate to provide thermal insulation between the die face and the extruder. The back plate preferably also functions as a heating plate for 95 which purpose it is preferably provided with a passage or passages for the circulation of a heating fluid: alternatively the back plate can embody or be surrounded by an electrical heater, but heating by means of a cir- 100 culating heating fluid is preferred. sired, a separate heating plate can be pro-vided between the back plate and the in-sulating plate. Each plate of the die-head is provided with feed channels, which are 105 in alignment when the plates are assembled together, through which the heat-plastified polymer is extruded and it has been found that any tendency for the heat-plastified polymer is solidified prematurely in these feed 110 channels with resulting impairment of the extrusion operation or even total blockage of the feed channels can be overcome in some circumstances (depending for example on the size of the feed channels and the extrusion temperature employed) by enlarging the diameter (or cross-sectional area) of the feed channels in the front plate constituting the die-face, or at least the extrusion orifice portions of such feed channels. ample the diameter of the extrusion orifices can be at least 20% greater than that of the feed channels in the back plate and insulating plate.

A convenient way of carrying out the present invention employing a face-cutting technique for manufacturing e.g. expandable polystyrene particles is illustrated in Figure II of the accompanying drawings. Referring to Figure II, polystyrene granules are fed 130

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to the hopper of an extruder 10 which is of the same kind as that described with reference to Figure I except for the die-head 35 which is of a sandwich construction as shown 5 on an enlarged scale in Figure III of the accompanying drawings and will be described in detail later with reference thereto. The pentane-containing polystyrene is extruded from the die-head and immediately 10 face-cut by a two-armed cutter 36 bearing against the face 37 of the die 35, whilst being cooled simultaneously to a temperature between 60°C and 80°C by water droplets sprayed against the die-face 37 from a nozzle 15 38 supplied with wet steam which condenses to form the water droplets in the vicinity of the die face 37. The resulting particles 39 are carried by the water droplets into slurry tank 40 containing expandable polystyrene particles in suspension in water at 60 to 80°C. A slurry of particles in water at this temperature is continuously pumped from the tank 40 to the base of a normalising tower 23 as described with reference to 25 Figure I; and the normalised particles are removed from the top of the normalising tower 23 and passed via a conveyor 30 to a storage bin 33, the arangement following the tank 40 being similar to that shown in Figure 30 I to which reference is made in this respect. The die-head, shown in Figure III, which is the preferred form of a die-head for use in the present invention when face-cutting is used, comprises a body of sandwich construc-35 tion mounted at the extrusion end of the ex-The die-head body comprises, in truder. combination, steel front and back plates 41, 42 with a cellulose-filled, phenol formaldehyde resin insulating plate 43 disposed be-40 tween the front and back plates 41, 42 to provide thermal insulation between the front plate 41, which provides the face portion 37 of the die-head, and the extruder 10. back plate 42, which is mounted on a flanged 45 portion 10' of the output end of the extruder 10, is provided with a plurality of feed channels 44 for the heat-plastified polymer. The back plate 42 is also provided with a milled channel 45 on its outer face which, when the 50 die-head is assembled and mounted on the extruder 10 with a plate 46 interposed between the back plate 42 and the extruder flanged portion 10, provides a passage through which steam under pressure is 55 passed to maintain the heating plate at about 100°C to 130°C depending on requirements. The plate 46 in addition to closing the channel 45 also provides transverse feed channels 47 in communication through orifices 48 with 60 the feed channels 44 in the back plate 42. The insulating plate 43 has a similar plurality of feed channels 49 each somewhat larger in diameter than the feed channels 44 in the black plate 42 and leading to a simi-65 lar number of feed channels 50 in the front

plate 41, the feed channels 50 being somewhat larger in diameter than the feed channels 49 in the insulating plate 43 and forming at their open ends 51 the extrusion orifices of the die-head.

The present invention is illustrated by the

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following Examples:— EXAMPLE I

Expandable polystyrene particles were manufactured by a strand-cutting method as 75 described above with reference to Figure I, the polystyrene being rendered expandable by the incorporation therein of technical pentane in a 60 millimetre (screw diameter) extruder comprising a heat-plastifying section 80 in which the polystyrene, supplied as granules to the feed hopper of the extruder, was lieated to 200°C; and a pentane-injection section maintained at 150°C to which liquid pentane was supplied and homogeneously in-corporated in the heat-plastified polystyrene prior to its extrusion into strands at the diehead at a temperature of 110°C. The resulting strands were immediately led into a water bath maintained at 65°C and, after a 90°C. short residence time therein, the strands were led out of the bath to an adjacent conventional strand cutter by which they were cut into particles 0.06 inch long and 0.02 inch in diameter (average dimensions). The particles immediately fell under gravity into a slurry tank maintained at 65°C from The 95 whence a slurry of particles was pumped to the base of a normalising tower in which normalising at 65°C was completed. The The 100 period in which the expandable polymer was out of contact with water at 60-80 °C, i.e. the time taken for the strand cutting operation, was of the order of 5 seconds. The residence time of the particles in the tower was 105 such that the total normalising period for the partieles was 40 minutes. After normalisation the particles were separated from the bulk of the water physically associated therewith and then air-dried.

The resulting expandable polystyrene particles contained on average 5% by weight of pentane and, when subjected to expansion conditions involving contact with live steam at ambient pressure, the particles expanded to form cylindrical beads, the lengths and diameters of which were similar and lay between 0.08 and 0.1 inches; and the beads had a cell size between 70 and 100 microns in diameter. The expanded particles were 120 found to be suitable in all respects for the manufacture of shaped articles e.g. blocks by conventional steam moulding technique. EXAMPLE II

Expandable polystyrene particles were 125 manufactured by a face-cutting method as described above with reference to Figures II and III, the polystyrene being rendered expandable by the incorporation of technical pentane in the manner described in Example 130

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I except that in order to avoid solidification of polymer in the die-head the temperature of the pentane-containing polystyrene at the die-head was 120°C.

A sprinkler was disposed adjacent to the front plate for spraying the face of the plate with water droplets at 70°C during operation of the extruder, the water droplets being formed by the condensation of wet steam.

10 The cutter was rotated at 30 revolutions per minute and the cut particles were carried down by the droplets of the water spray into a body of water, also at 70°C, and after a residence time therein of 40 minutes to

15 achieve normalisation, the particles were separated from the water and cooled to ambient temperature and simultaneously dried

in a current of air.

Using polystyrene and operating at the 20 specified extrusion temperature of 120°C, particles of about 1 millimetre in diameter and containing 5% by weight of pentane were obtained which showed no sign of When subjected to expansion con-25 ditions involving contact with live steam at ambient pressure the particles expanded to form generally spherical beads having a diameter between 3 and 5 millimetres and a cell size ranging from 70 to 100 microns in dia-30 meter which were found suitable in all respects for the manufacture of shaped articles by conventional steam moulding technique. **EXAMPLE III**

Expandable polystyrene particles were 35 manufactured by the face-cutting method the die-head being a multi-orifice die-head from which 32 strands of pentane-containing polystyrene were extruded simultaneously, the

die-head comprising a body of sandwich con-40 struction having a face portion against which a bladed cutter, rotated during operation at 30 revolutions per minute, was disposed in such a manner as to cut the extrudate into particles at the die face. The die-head body

45 comprised, in combination, steel front and back plates with a cellulose-filled phenol formaldehyde resin insulating plate disposed between the front and back plates to provide heat insulation between the front plate,

50 which formed the face portion of the diehead, and the extruder. The back plate, which was mounted on the extruder at its output end, was provided with 32 feed channels each 1.5 millimetres in diameter.

55 insulating plate was 1.5 millimetres thick and had 32 feed channels each 3 millimetres in diameter and the front plate, which was 2.5 millimetres thick, also had 32 feed channels of the same diameter as those in the in-

60 sulating plate, the feed channels in the front plate and in the insulating plate being in register with one another and with those in the back plate. The die-head also embodied a separate heating plate which comprised 65 a copper plate having 32 feed channels each 1.5 millimetres in diameter and was mounted between the insulating plate and the back The heating plate had a thickness of 6 millimetres and was provided with a spiral milling on its outerface which, when the die- 70 head was assembled and mounted on the extruder, provided a 3 millimetre-square channel through which steam was passed to maintain the heating plate at about 100°C.

The extruder employed comprised a 60 75 millimetre (screw diameter) extruder fitted with a screw having a 20:1 length to diameter ratio and a compression ratio of 3:1 in the heat-plastifying section, the screw being of a design providing a heat-plastifying 80 section in which rapid compression over an initial, one-diameter length of the screw took place; followed by a five-diameter length dispersion section to which technical pentane was supplied for incorporation in the heat- 85 plastified polystyrene. In operation the screw was rotated at 20 revolutions per minute and the temperature conditions in the extruder were such that the temperature profile thereof increased from 80°C at the input 90 end to 100° C at the die-head end, the actual temperature of the pentane-containing heatplastified polystyrene during extrusion being

110°C.

The face-cutting operation was carried out 95 under a spray of water droplets at 60°C, the cut particles being carried down by the droplets of the water spray into a body of water. also at 60°C, and after a residence time therein of 1 hour to achieve normalisation, 100 the particles were separated from the water and cooled to ambient temperature and simultaneously dried in a cufrent of air. When operating at the specified extrusion temperature of 110°C, particles of about 3 milli- 105 metres in diameter and containing 4.8% by weight of pentane were obtained which showed no sign of voids. On expansion with live steam at ambient pressure such expandable polystyrene particles expanded to 110 form generally spherical beads having a diameter of about I centimetre and a cell size ranging from 80 to 100 microns in diameter. Such particles were moulded into blocks of cellular polystyrene by conventional steam 115 moulding technique and performed satisfactorily in this application.
WHAT WE CLAIM IS:-

1. A process for the manufacture of expandable particles of a thermoplastic organic 120 polymer (as hereinbefore defined) in which a varporisable expanding agent is incorporated in a heat-plastified mass of said polymer and the heat-plastified polymer is extruded in filamentary form and cut into par- 125 ticles, which comprises extruding the heatplastified polymer, containing said expanding agent, into an essentially gaseous environment at atmospheric or substantially atmospheric pressure and immediately cool- 130

ing the extruded polymer by contact with a fluid at a temperature between 60 and 80°C in such a manner that the temperature of the extruded polymer does not fall below 60°C, 5 maintaining the temperature of the extruded polymer between 60 and 80°C during a normalising period (as herein defined) immediately following extrusion thereof and before and/or after cutting the extruded polymer 10 into particles, and thereafter cooling the polymer particles to ambient temperature.

2. A process as claimed in claim 1, which comprises cutting the extruded polymer into particles whilst cooling said extruded poly15 mer by contact with said fluid at a tempera-

ture between 60 and 80°C.

3. A process as claimed in claim 2, wherein the polymer is extruded from a dichead at which the extruded polymer is face20 cut into particles whilst being simultaneously cooled to a temperature between 60 and 80°C by droplets of a liquid sprayed against the face of said die-head, and the resulting particles are then immediately con25 tacted with a body of said liquid at 60-80°C.

4. A process as claimed in claim 3, wherein the die-head comprises an outer face plate and an insulating plate providing thermal insulation between said face plate and 30 the body of the extruder to which said die-

head is attached.

t5. A process as claimed in claim 4, wherein said die-head comprises a heating plate interposed between said insulating
plate and said extruder, said heating plate comprising a passage or passages for the circulation of a fluid at elevated temperature.

6. A process as claimed in claim 1, which comprises immediately contacting the ex-40 truded polymer with a body of fluid at a temperature between 60 and 80°C, cutting the filamentary extruded polymer into particles and substantially immediately contacting the resulting particles with said fluid at 45 60-80°C.

45 60-80°C.

7. A process as claimed in claim 6, wherein the polymer is extruded into a body of liquid at 60-80°C from a die-head located adjacent to but spaced from said 50 body of liquid and the resulting filamentary extruded polymer is removed from said body of liquid for cutting into particles, which particles are then immediately contacted with a body of liquid at 60-80°C.

8. A process as claimed in any one of the preceding claims, wherein said thermoplastic organic polymer is polystyrene. 9. A process as claimed in any one of the preceding claims, wherein said vaporisable expanding agent consists of or comprises u- 60 pentane or iso-pentane or a mixture thereof.

10. A process as claimed in any one of the preceding claims, wherein the extrusion temperature of said heat-plastified polymer is between 130 and 160°C and the polymer 65 is extruded through orifices below 1.0 millimetre in diameter.

11. A process as claimed in any one of the preceding claims, wherein said fluid or

liquid at 60-80°C is water.

12. A process as claimed in any one of the preceding claims, wherein at least the major part of the normalisation of the extruded polymer is carried out in a tower containing water maintained at 60 to 80°C, a suspension of polymer particles in water at 60 to 80°C being supplied continuously to the base of said tower and said particles being moved progressively up said tower for discharge from the top of said tower.

13. A process for the manufacture of expandable particles of a thermoplastic polymer substantially as hereinbefore described with reference to Figure I of the accompany-

ing drawings.

14. A process for the manufacture of expandable polystyrene particles substantially as hereinbefore described with reference to Example I.

15. A process for the manufacture of expandable particles of a thermoplastic polymer substantially as hereinbefore described with reference to Figures II and III of the accompanying drawings.

16. A process for the manufacture of expandable polystyrene particles substantially as hereinbefore described with reference to

Example II or Example III.

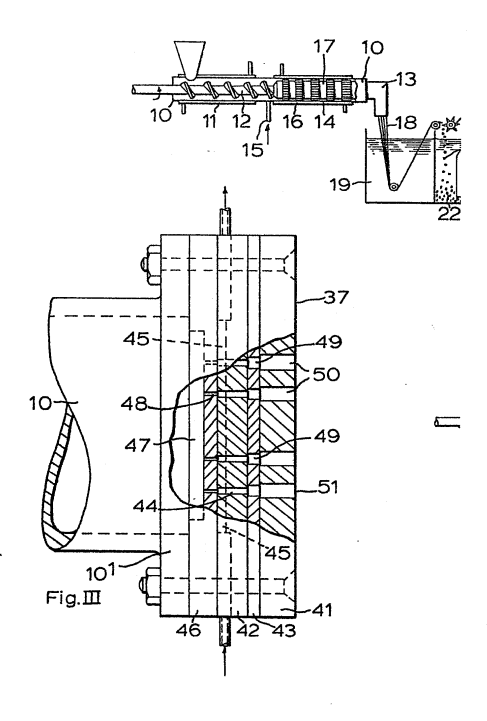
17. Expandable particles of a thermoplastic polymer when manufactured by the 100 process claimed in any one of the preceding

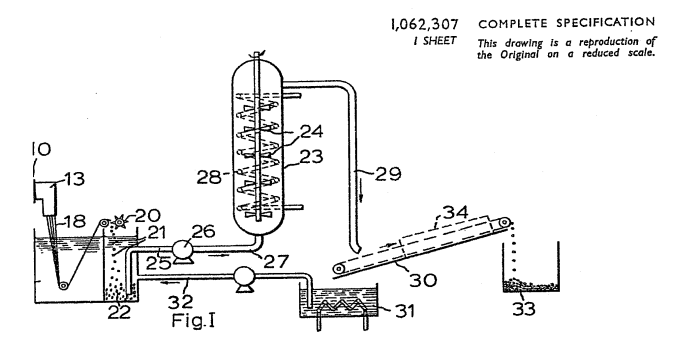
18. Expanded particles of a thermoplastic polymer obtained by subjecting to expansion conditions expandable particles as claimed 105 in claim 17.

19. A shaped article manufactured by moulding expanded particles as claimed in claim 18.

WILLENS & ROBBINS
Chartered Patent Agents
Shell Centre
London, S.E.1.
Agents for the Applicants.

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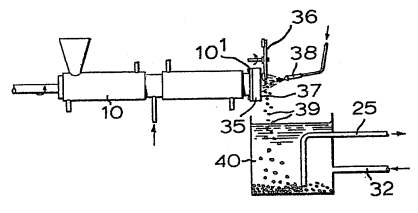


Fig. I

